## Amendments to the Claims:

- 1 5. (Canceled)
- (Currently amended) A medical device for implanting in a patient, comprising: a device body with <u>a an-outer surface</u>;

an attachment region within the surface, wherein the attachment region comprises a cavity in the surface having an open end and an opposing closed end, the cavity having a base surface at the closed end and a side wall extending from the closed end to the open end; and a ceramic component comprising

a first porous <u>ceramic</u> region, and
a second porous <u>ceramic</u> region, wherein the second porous <u>ceramic</u>
region is less porous than the first porous <u>ceramic</u> region, the ceramic
component connects to the attachment region through the second
porous <u>ceramic</u> region <u>coupled to the base surface of the closed end</u>, and
the second porous <u>ceramic</u> region is positioned in between the first porous
<u>ceramic</u> region and the <u>base surface of the</u> attachment region
such that the
first porous <u>ceramic</u> region and <u>the base surface of the</u> attachment region
are located on opposite sides of the second porous ceramic region:

wherein the attachment region comprises an indentation in the surface
wherein the device is a stent.

- (Previously Presented) The medical device of claim 6 wherein one or both of the porous regions releasably contains a drug.
- 8. (Original) The medical device of claim 7 wherein the drug comprises at least one of a smooth-muscle-cell vascular activity inhibitor, a wound healing enhancer, an agent for improving the structural properties in a vascular site, an agent for improving the elastic properties of a vascular site, an antineoplastic substance, an anti-inflammatory substance, an antiplatelet substance, an antitoagulant substance, an antifibrin substance, an antithrombin

substance, an antimitotic substance, an antibiotic substance, an antiallergy substance, an antioxidant substance, alpha-interferon, genetically engineered epithelial cells, rapamycin, actinomycin D, paclitaxel or docetaxel.

- (Withdrawn) The medical device of claim 6 further comprising a polymer layer over the ceramic component, over a portion of the medical device not including the ceramic component, or both.
- 10. (Withdrawn) The medical device of claim 6 further comprising an auxiliary component with at least one auxiliary-component attachment region disposed in or on the surface of the auxiliary component and wherein the ceramic component is disposed on or within at least one auxiliary-component attachment region.
- 11. (Withdrawn) The medical device of claim 10 further comprising a third porous region disposed in the ceramic component wherein the third porous region is less porous than the first and wherein the ceramic component connects to at least one auxiliary-component attachment region through the third porous region.
- (Withdrawn) The medical device of claim 11 wherein the ceramic component is fused to at least one auxiliary-component attachment region through the third porous region.
- (Withdrawn) The medical device of claim 11 further comprising an oxide layer disposed between the third porous region and at least one auxiliary-component attachment region.
- (Withdrawn) The medical device of claim 11 wherein the surface or auxiliarycomponent surface, or both, comprise a metal, glass, or ceramic.
- 15. (Withdrawn) The medical device of claim 14 wherein metal comprises iron, cobalt, nickel, manganese, stainless steel, tantalum, niobium, super-elastic nickel-titanium alloys, titanium, silver, gold, platinum, steel, or aluminum.
- (Withdrawn) The medical device of claim 14 wherein glass comprises borosilicate glass, lead glass, soda glass, uranium glass, soft glass, fused quartz, or fused silica.

- (Withdrawn) The medical device of claim 14 wherein ceramic comprises carbide ceramics, oxide ceramics, nitride ceramics, or boride ceramics.
- 18. (Withdrawn) The medical device of claim 14 wherein ceramic comprises titania, zirconia, hafnia, silica, alumina, silica alumina, silicon carbide, tungsten carbide, silicon boronitride, boronitride, silicon, or gallium arsenide.
- (Withdrawn) The medical device of claim 10 wherein the auxiliary component is one of an electrode, a physical sensor, or a chemical sensor.
- 20. (Withdrawn) The medical device of claim 10 further comprising a polymer layer disposed over the auxiliary component, over a portion of the medical device not including the auxiliary component, or both.

## 21. (Canceled)

- (Previously presented) The medical device of claim 6 wherein the surface of the medical device comprises plastic, metal, glass, or ceramic.
- (Original) The medical device of claim 22 wherein metal comprises iron, cobalt, nickel, manganese, stainless steel, tantalum, niobium, super-elastic nickel-titanium alloys, titanium, silver, gold, platinum, steel, or aluminum.
- (Withdrawn) The medical device of claim 22 wherein glass comprises borosilicate glass, lead glass, soda glass, uranium glass, soft glass, fused quartz, or fused silica.
- 25. (Withdrawn) The medical device of claim 22 wherein ceramic comprises carbide ceramics, oxide ceramics, nitride ceramics, or boride ceramics.
- 26. (Withdrawn) The medical device of claim 22 wherein ceramic comprises titania, zirconia, hafnia, silica, alumina, silica alumina, silicon carbide, tungsten carbide, silicon boronitride, boronitride, silicon, or gallium arsenide.

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Appl. No. 10/623,908 Amendment dated April 24, 2009 Reply to Office Action of November 24, 2008

(Currently Amended) A medical device for implanting in a patient comprising:
 a) a surface comprising a metal;

b) an attachment region disposed within the surface, wherein the attachment region comprises an indentation in the surface;

c) a ceramic component comprising a glass or ceramic, the ceramic component having a first porous <u>ceramic or glass</u> side and a second less porous <u>ceramic or glass</u> side, wherein the less porous <u>ceramic or glass</u> side of the ceramic component is fused on or within the attachment region; and

d) an oxide layer disposed on or within the attachment region between the surface of the device and the ceramic component;

## wherein the medical device is a stent.

- (Canceled)
- (Previously Presented) The medical device of Claim 27 further comprising a drug releasably disposed in the first porous side.
  - 30. 46. (Canceled)
- 47. (Currently amended) The medical device of claim 6 wherein an oxide layer is disposed between the attachment region and the second porous <u>ceramic</u> region.
- 48. (Previously presented) The medical device of claim 47 wherein the oxide layer comprises an oxide of the material of which the medical device body is comprised.
- (Previously presented) The medical device of claim 6 wherein the attachment region is created by removing some of the material from the medical device body.
- (Previously presented) The medical device of claim 6 wherein the ceramic component comprises a compound selected from the group consisting of carbide ceramics, oxide ceramics, nitride ceramics, boride ceramics, and combinations thereof.

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- 51. (Previously presented) The medical device of claim 6 wherein the ceramic component comprises a compound selected from the group consisting of borosilicate glass, lead glass, soda glass, uranium glass, soft glass, fused quartz, fused silica, and combinations thereof.
- (Currently amended) The medical device of claim 6 wherein the surface of the attachment region is machined to more closely matches the thermal characteristics of the ceramic component.